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Des ription

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FIELD OF THE INVENTION

The present invention relates to a resin composition having a prolonged shelf-life which is capable of providing a cured product having good thermal resistance and good impact strength, and also relates to a process for the production of the composition. More particularly, the present invention relates to a resin composition which gives a molded product having both the excellent mechanical and thermal characteristics inherent to an epoxy resin and the excellent toughness inherent to a thermoplastic resin.

BACKGROUND OF THE INVENTION

Recently, a composite in which carbon fibers, aromatic polyamide fibers or the like are used as reinforcement have been frequently used as structural materials for aircraft and the like, based on their high specific strength and high specific rigidity.

Composites which have excellent thermal resistance, mechanical characteristics, dimensional stability, chemical resistance and weather resistance have been provided by combining aromatic glycidylamine epoxy resins as the matrix resin with a diaminodiphenylsulfone as a curing agent.

Though such composites prepared from epoxy resin-based prepregs show good performance, it has been found that they have inferior toughness and impact strength because of the short shelf-life of the prepreg and the low elongation and, therefore, brittleness of the matrix resin.

Especially, an improvement of the impact strength of the composite without reducing the thermal resistance thereof has been regarded as an important object in the case of the use of such composites as primary structural materials in aircraft which are apt to receive impact from the outside, for instance, by pebbles bounced up at the time of take-off or landing, from tools dropped by mistake during maintenance and repair and the like.

In order to increase the high impact strength of a composite, it is important not only to improve the elongation of a reinforcement such as carbon fibers but also to increase the toughness of a matrix resin. For this reason, a number of attempts have been made to improve matrix resins.

In order to improve the toughness of matrix resins, several methods have been proposed such as a method in which an epoxy resin is mixed with a rubber component and a method in which an epoxy resin is mixed with a high molecular weight component. When an epoxy resin is mixed with a rubber component, the toughness and impact strength of the resulting molded product can be improved, but its thermal resistance and mechanical characteristics are reduced. Because of this, the mixing ratio of the rubber component is limited, sometimes to a low mixing ratio depending on the use of the product, thus resulting in insufficient improvement.

In order to mix an epoxy resin with a thermoplastic resin as a high molecular weight component, a thermoplastic resin is dissolved in the epoxy resin at high temperature, or a thermoplastic resin is dissolved in a solvent and then the epoxy resin is added to the solution. When a thermoplastic resin is dissolved in an epoxy resin at a high temperature, the viscosity of the resin mixture increases and, at the same time, its tackiness decreases, thus resulting in markedly poor handling ability. Mixing of these resins using a solvent also has disadvantages such as the problem of removing the solvent after mixing, complex preparation steps and a decrease in thermal resistance due to a small amount of remaining solvent.

As a consequence, conventional resin compositions prepared by mixing an epoxy resin with a small amount of a rubber component or a high molecular weight component provide a composite with a poor impact strength-improving effect though the decrease in thermal resistance is not so great.

Other types of resin compositions used as prepregs have been disclosed, for instance, in JP-A-61-250032, JP-A-62-57417 and JP-A-63-162732 (corresponding to U.S. Patent 5,028,478), in which a thermoplastic resin is dispersed in and mixed with an epoxy resin to improve toughness (impact strength) of a composite, but with insufficient effect in terms of impact strength of the composite. (The term "JP-A" as used herein means an "unexamined published Japanese Patent Application".)

In addition, in these resin compositions, a diaminodiphenylsulfone compound is used as an epoxy resincuring agent in view of the thermal resistance of the resulting composite. When such a compound is used as a curing agent, the resulting resin composition has a short shelf-life of about 2 to 3 weeks at room temperature (about 23 °C), thus resulting in a handling problem.

EP-A-0 392 348 discloses a resin composition which comprises an epoxy resin; a thermoplastic resin; and particles of a diaminodiphenylsulfone (DDS) compound.

CA-A-716 638 discloses a curing agent which can be pre-mixed with polyepoxid s, and wher in the mixture can be stor d without curing. The curing agent may b formed of small particles of DDS whose reactivity at room t mperatur is inhibited by a hardened protective coating.

Accordingly, this invention contemplates over-coming th abov -m ntion d problems involved in the prior art.

SUMMARY OF THE INVENTION

The first object of the present invention is to provide a resin composition which has a prolonged shelflife and is capable of providing a composite having excellent thermal resistance and excellent mechanical characteristics such as impact strength, toughness and the like, and to provide a process for the production of such a composition.

The second object of the present invention is to provide a mixed type resin composition having advantages inherant to both of an epoxy resin and a thermoplastic resin and to provide a process for the production of such a composition.

The third object of the present invention is to provide a resin composition suitably used as a prepreg and to provide a process for the production of such a composition.

The resin composition of the present invention comprises:

(A) an epoxy resin,

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- (B) a thermoplastic resin selected from the group consisting of polyether imide, polyether sulfone, polysulfone, polycarbonate, polyetherether ketone, polyamide, and a mixture of two or more of these;
 - (C) particles of a diaminodiphenylsulfone compound whose cross-linking reactivity with the epoxy resin is prevented until the resin composition is subjected to molding or curing (i) by resin particles adhered on the surface of the diaminodiphenylsulfone particles or (ii) by a resin film coated thereon; the resin forming said resin particles or resin film being a polyamide, a modified urea resin, a cured melamine resin, a polyolefin, a paraffin, a modified paraffin or a mixture of two or more of these.

The resin composition of the present invention can be obtained by a method which comprises mixing: (A) an epoxy resin,

- (B) a thermoplastic resin selected from the group consisting of polyether imide, polyether sulfone, polysulfone, polycarbonate, polyetherether ketone, polyamide, and a mixture of two or more of these; and
- (C) particles of a diaminodiphenylsulfone compound whose cross-linking reactivity with the epoxy resin is prevented until the resin composition is subjected to molding or curing (i) by resin particles adhered on the surface of the diaminodiphenylsulfone particles or (ii) by a resin film coated thereon; the resin forming said resin particles or resin film being a polyamide, a modified urea resin, a cured melamine resin, a polyolefin, a paraffin, a modified paraffin or a mixture of two or more of these.

Other objects and advantages of the present invention will be made apparent as the description progresses.

DETAILED DESCRIPTION OF THE INVENTION

Examples of an epoxy resin which can be used as component (A) in the present invention include glycidylamine epoxy resins, bisphenol epoxy resins, novolak epoxy resins, urethane-modified bisphenol A epoxy resins, alicyclic epoxy resins, and the like. The combined use of these resins with a diaminodiphenylsulfone compound as a curing agent makes possible the formation of a molded product having high thermal resistance and excellent mechanical characteristics.

Specific examples of each type of epoxy resins are recited below:

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(1) Glycidylamine Epoxy Resin

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(2) Bisphenol Epoxy Resin

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Bisphenol A Epoxy Resin

n = 1-34Bisphenol F Epoxy Resin (x : = CH₂) n = 1

Bisphenol AD Epoxy Resin

n = 1 Bisphenol S Epoxy Resin (x : = SC₂) n = 1

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- (3) Novolak Epoxy Resin
- i) Phenol Novolak Epoxy R sin

O-CH₂-CH-CH₂
O-CH₂

n > 0

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ii) Cresol Novolak Epoxy Resin

CH₃ O-R CH₂-CH-CH₂

CH₃ O-CH₂-CH-CH₂

CH₂

CH₂

CH₃ O-CH₂-CH-CH₂

R: H or CH₃ n ≥ 0

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(4) Urethane Modified Epoxy Resin

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Bisphenol A poxy resin modified with urethan at both ends thereof.

5	Ţ	â				
10		Tradename of Product Produced by CIBA-GEIGY u, c, c,	ERL4234	ERL4299	ERL4221	ERL4206
15		ne of Pro EIGY	.75	121	CY179	
20		Tradename of CIBA-GEIGY	CX175	CX177	Ċ.K.	
25		Viscosity	125,000	550-750	350-450	<15
30	,	Epoxy Equivaleent Viscosity (qr/mol epoxy)	154	.210	131-143	70-74
35		Epoxy E	133-154	190-210	131-	70.
40	esin) acetal	H274C-O-CH2	CH2-O-C O Alicyclic diepoxy carboxylate	dioxide
45	Epoxy Re		diepory	cłch ₂ h ₄ c	C.C.	H ₂
50	(5) Aliphatic Epoxy Resin		OT Cyclic diepoxy acetal	CH2-O-C+CH2)4C-O-CH2	CH2-0-	Vinyl cyclohexene dioxide
	Ü	·			<u> </u>	<u></u>

Commercially available epoxy resins are shown below: Examples of glycidylamine epoxy resins include MY-720 (Ciba-Geigy Ltd.), Epotohto YH 434 (Tohto

Kasei Co., Ltd.), Ep. 604 (Yuka Sh II Co., Ltd.), ELM-120 (Sumitomo Ch mical Co., Ltd.), ELM-100 (Sumitomo Ch mical Co., Ltd.), GAN (Nippon Kayaku Co., Ltd.) and th lik .

Exampl s of bisphenol epoxy r sins includ Epikote 815, Epikote 828, Epikote 834, Epikote 1001 and Epikote 807 (Yuka Shell Co., Ltd.), Epomic R-170 (Mitsui Petrochemical Industri s, Ltd.), EXA 1514 (Dainippon Ink & Chemicals, Inc.) and the like.

Examples of novolak epoxy resins include phenol novolak epoxy resins such as Epikote 152 and Epikote 154 (Yuka Shell Co., Ltd.), Dow Chemical DEN 431, DEN 485 and DEN 438 (Dow Chemical Co.), Epikote 154 (Yuka Shell Co., Ltd.), Dow Chemical DEN 431, DEN 485 and DEN 438 (Dow Chemical Co.), Epikote 154 (Yuka Shell Co., Ltd.) and the like, and cresol novolak epoxy resins such as Epikote 152 and ECN 1240 (Ciba-Geigy Ltd.), EOCN 102, EOCN 103 and EOCN 104 (Nippon Kayaku Co., Ltd.) and the like.

Alicyclic epoxy resins such as Araldite CY-179, CY-178, CY-182 and CY-183 (Ciba-Geigy Ltd.) and the like are also useful.

As urethane-modified bisphenol A epoxy resins, Adeka Resin EPU-6 and EPU-4 (Asahi Denka Kogyo K.K.) and the like are commercially available. When these resins are used, a resin composition having excellent flexibility and good adhesion to reinforcing fibers can be obtained. In addition, since these resins have the function of creating a heterogeneous structure of the resin due to their poor compatibility to the thermoplastic resin, they can contribute to the improvement of impact strength.

A reactive diluent may also be incorporated into the resin composition of the present invention as a part of component (A), if desired. The amount of the diluent is preferably not more than 10% by weight based on the weight of component (A).

Examples of the diluent include polypropylene diglycol diglycidyl ether, phenylglycidyl ether or the like. Epikote 871 or Epikote 872 (dimer acid modified epoxy resin; Yuka Shell Co., Ltd.), TACTIX 695 (a mixture comprising bisphenol A and brominated bisphenol A; Dow Chemical Co.) or the like may also be added as a flexible epoxy resin. Also useful as an additive is a heat resistant epoxy resin such as Epikote 1031 (tetrafunctional epoxy resin) or Epikote 1032 (hepta-functional epoxy resin) (Yuka Shell Co., Ltd.), TACTIX 742 (tri-functional epoxy resin; Dow Chemical Co.) or the like.

Typical examples of the thermoplastic resin as component (B) in the present invention include polyether imide, polyether sulfone, polysulfone, polyetherether ketone, polyamide, and the like, of which polyether imide, polyether sulfone and polysulfone are particularly preferred because of their excellent thermal resistance.

Specific examples of each type of thermoplastic resins are shown below:

(1) Polyether Imide

(2) Polyether Sulfone

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(3) Polysulfone

(4) Polycarbonate

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(e.g., n = 140)

(5) Polyetherether Ketone

35 (6) Polyamide

wherein R and R' each represents an alkylene or an arylene group, preferably having 1 to 12 and 6 to 12 carbon atoms, respectively.

These thermoplastic resins may be used alone or as a mixture of two or more of them, and these resins may be in a form of particles from the point of view of preparation of the resin composition of the present invention. A preferred size ranging from 1 to 50 μ m, preferably from 1 to 20 μ m.

Component (B) is preferably used in an amount of from 5 to 40 parts by weight, more preferably of from 15 to 35 parts by weight per 100 parts by weight of component (A).

Amounts of component (B) if larger than 40 parts by weight based on 100 parts by weight of an epoxy resin as component (A) usually tend to result in a high resin viscosity which leads to difficulty in attaining uniform mixing. Amounts of component (B) if smaller than 5 parts by weight tend to reduce toughness and, th refore, improvem nt in th impact strength also tends to decr ase.

The diaminodiph nylsulfone compound (hereinafter abbr viated as DDS) used in the present invention can be represented by the following formula:

$$\mathsf{NH}_2 \\ \mathsf{SO}_2 \\ \mathsf{NH}_2$$

Examples of the DDS include 3,3'-diaminodiphenylsulfone, 3,4'-diaminodiphenylsulfone and 4,4'-diaminodiphenylsulfone. The DDS is used as fine particles having a size preferably of from 1 to 50 μ m, and more preferably from 10 to 30 μ m.

In the present invention, in order to prolong or extend the shelf-life of the composition which is a mixture of components (A), (B) and DDS or is in a form of prepreg thereof, the activity of the cross linking reactivity (or curing function) of the DDS is prevented or protected until the composition is subjected to molding or curing.

In the present invention DDS should be prevented from reacting with the epoxy resin tridimensionally to form a cured epoxy resin by cross linking. Proceeding of a unidimensional reaction (by which a tridimensional molecular structure as in the molecules of a cured epoxy resin does not form) of the DDS and the epoxy resin to form a linear compound is permitted.

Such desired effect can be attained by adhering fine resin particles on the surface of each DDS particle. The resin particles may be adhered to the surface separately from each other, may form a network structure, or may form a layer on the surface of the DDS particle, which layer may be continuous or non-continuous, or porous or non-porous, so long as the molten DDS can flow out through the spaces among the resin particles, through the pores or through the non-continuous portion of the layer, and/or the DDS particles can be exposed due to melting or destruction of the covering comprising the resin particles, under the conditions (pressure and/or temperature) of molding or curing.

In the present invention DDS particles are coated with a resin having less or no reactivity to epoxy resin, in order to prevent reaction of the DDS with an epoxy resin.

The size of the resin particles is less than the size of the DDS particles. A preferred particle size of the resin particles is from 1/3 to 1/100 of that of the DDS particles, and it is usually from 0.1 to 5 μ m.

The molding of the composition is usually conducted at a temperature of from about 90 to 200 °C and a pressure of from about 1 to 5 kgf/cm². The curing temperature is usually from about 170 to 200 °C. The resin which can be used for coating may be selected from those which melt or of which covering is destroyed at the molding temperature and/or under the molding pressure, or at the curing temperature.

The resin used for the coating of the DDS particles is selected from thermoplastic resins or thermosetting resins. Illustrative examples of resins which can be used for coating in the present invention include polyamide resin, modified urea resin, cured melamine resin, polyolefin, polyparaffin (including modified products) and the like.

When the resin covering has a structure through which the molten DDS is able to flow out, the type of the resin can be selected from resins which do not melt or those of which the covering may not be destroyed under the above-described conditions.

These coating resins may be used alone or as a mixture of two or more of them. Alternatively, two or more types of DDS particles which are coated using different coating resins may be used in the resin composition of the present invention.

Adhesion of particles on the surface of the DDS particle can be carried out by conducting a solution polymerization in a liquid dispersion of the DDS particles so that polymer particles are formed on the surface of the DDS particles, or by curing a resin in a resin solution or emulsion wherein the DDS particles are dispersed to form cured resin particles on the surface of the DDS particles. The method which is disclosed in detail in JP-A-3-4932 and JP-A-3-238030 can be preferably used for production of component (C) of the present invention. In this method DDS particles and an amino compound/formaldelhyde primary polycondensation product are dispersed in an aqueous solution of a water soluble polymer to form an emusion. By heating the emusion while stirring the polycondensation reaction is proceeded to deposite the polycondensation product on the surface of the DDS particles as fine particles to coat the surface of the DDS surface. Further, adhesion of the resin particles or forming of a resin film on the surface of the DDS particles can be carried out by a dry method in which the DDS particles and the resin particles for adhering thereto or for forming film thereon are mixed using a high speed mixer or the like to adhere the particles of coating resin lextrostatically on the surface of the DDS particles, and then the resin particles are firmly adhered on the surface of DDS particles by heat-melting the resin or the resin particles are made into a film form by heat-melting the resin. These methods may b selected properly dep nding on the properties of the coating resin and the particle size desired.

In order to gain good composite characteristics, it is desirable to coat the coating resin uniformly and thinly on the surface of the DDS particles. For this purpose, the coating resin may pr ferably b used in an amount ranging from 5 to 20% by weight, more preferably from 10 to 15% by weight based on the weight of the DDS particles.

Since reaction of the epoxy resin with the curing agent hardly or does not progress at room temperature, the resin composition of the present invention has a longer shelf-life (about twice) that of a resin composition in which DDS is used as a curing agent without coating it by a coating resin. When the resin composition of the present invention is subjected to molding or curing, the DDS compound starts to react with the epoxy resin when the temperature reaches about 170 °C or higher, thereby resulting in the formation of a cured product.

Other curing agents or curing accelarators may be added to the composition in such amounts that the shelf-life of the composition and the physical properties of the composite obtained therefrom are not altered by the addition of such agents. It is preferred that the amount of such agents or accelerator is not more than 10 parts by weight per 100 parts by weight of component (A).

Examples of such epoxy resin-curing agents include aromatic amines such as metaphenylenediamine, diaminodiphenylmethane and the like; acid anhydrides such as phthalic anhydride, trimellitic anhydride, pyromellitic anhydride and the like; boron trifluoride complex salts such as BF₃ monoethylamine, BF₃ benzylamine and the like; and imidazoles such as 2-ethyl-4-methyl imidazole, 2-ethyl imidazole, 2,4-dimethyl imidazole, 2-phenyl imidazole and the like. Urea compounds (3-[3,4-dichlorophenyl]-1,1-dimethyl urea and the like) and organic metal salts (Co(III) acetyl acetonate and the like) may also be used jointly.

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The DDS as an epoxy resin-curing agent is used in an epoxy equivalent/amine equivalent ratio of 1/1 in theory but, taking the mechanical properties and the water absorption coefficiency of the resulting cured product into consideration, the DDS may be used generally within the equivalent ratio of 1/(0.6 to 1.3), preferably 1/(0.8 to 1).

In addition to the aforementioned essential components, the resin composition of the present invention may further contain rubber components (for example, a carboxyl-terminated butadiene/acrylonitrile copolymer, nitrile rubber, epoxy-modified polybutadiene rubber and the like) in small amounts so that the thermal resistance of the composition is not reduced and fillers (silica powder and the like for example) in such amounts that handling of the resulting prepreg does not worsen, as well as flame retardants such as antimony trioxide, coloring agents and the like. From a handling point of view, the resin composition of the present invention may also contain small amounts of a modified acrylic polymer (MODAFLOW manufactured by Monsanto Co., for example) as a flowability regulating agent and silicone resin, silicone oil, vaseline or the like as a water repellent. These substance are added in a total amount preferably of not more than 10 parts by weight per 100 parts by weight of component (A).

The resin composition of the present invention may be prepared, for instance, in accordance with the following illustrative procedure.

Each of the components is sudjected to heat kneading using a kneading machine preferably in an inert gas atmosphere. The heating temperature in this instance is set to a level lower than the curing initiation temperature of the epoxy resin used. In general, the ingredients of component (A) are firstly mixed uniformly at a temperature of from 20 to 90 °C when component (A) comprises two or more epoxy resins, preferably from 40 to 90 °C. Component (A) is mixed with components (B) and (C) to form a uniform mixture at a temperature of from 20 to 90 °C. In this instance, component (B) may be used in a finely powdered form preferably having a particle size of from 1 to 50 μm, more preferably from 1 to 20 μm, so that component (B) can be dispersed uniformly and preparation of the resin composition can be carried out smoothly. In this manner, a resin composition can be prepared in which a thermoplastic resin component is blended to a high level of about 40 parts by weight per 100 parts by weight of the epoxy resin. However, blending of the thermoplastic resin component to a higher level than 40 parts by weight usually is not preferable because the viscosity of the composition becomes too high which leads to difficulty in kneading the components.

The resin composition of the present invention is suitable for use to produce a molded product, especially, containing fibers as reinforcement.

Examples of fiber reinforcement which are preferably used in the present invention include carbon fiber, glass fiber, aromatic polyamide fiber and the like, which may be used alone or in combination. For the purpose of improving the mechanical properties of a composite, it is preferable to use so-called "high strength medium modulus of elasticity carbon fibers" which have a t nsile strength of 400 kgf/mm² or more and a modulus of elasticity of 30×10^3 kgf/mm² or more. The modulus of elasticity is preferably not more than 40×10^3 Kgf/mm².

The length of the carbon fiber usually is from 5 to 70 mm, or the carbon fiber may be continuous. The fiber reinforcement may be in the form of a textile, a unidirectional parallel yarn product, chopped fiber or the like. The fiber reinforcement has a diameter of from 1 to 20 μ m, preferably from 5 to 7 μ m. The preferable amount of the fiber reinforcement is from 5 to 70% by volume, and preferably from 55 to 65% by volume, based on the volume of the composition. Carbon fibers which are coated with a metal such as Ni, Al, Cu Zn, Pb, Su, Au, Ag may also be used in the present invention.

In the process of the present invention, a prepreg for production of a molding is generally prepared by impregnating the resin composition of the present invention into a fiber reinforcement, which preferably is effected by a conventional hot melt method. This method is carried out by preparing a resin composition film, placing fibers thereon, and hot-pressing the thus obtained assembly. This method is described more in detail, for example, in U.S. Patents 4,714,648, 4,482,660, and 4,309,473.

The prepreg containing the fiber obtained by such a process has excellent qualities. The prepreg is subjected to molding and curing, usually using a hot-press method or an autoclave method. The hot-press method is usually conducted at a temperature of from about 90 to 200 °C and a pressure of from about 1 to 5 kgf/cm² for about 1 to 3 hours.

Since the resin composition of the present invention is preferably prepared by uniformly dispersing the component (B) thermoplastic resin in the component (A) epoxy resin, not only it is possible to prepare a prepreg by the conventional hot melt method, but also it is possible to prepare an excellent resin composition for a prepreg having a prolonged shelf life. The product obtained from the prepreg does not raise problems due to remaining solvent and has both the excellent thermal stability inherent to the epoxy resin and the toughness and impact strength inherent to the thermoplastic resin without decreasing the mechanical characteristics thereof.

Though the reason is not known, a composite having excellent toughness and impact strength can be obtained when DDS particles are coated and used as a curing agent of a resin composition comprising components (A) and (B). Such effects are superior to the case of a separate mixing of the DDS particles with the coating resin without coating it on the DDS particles.

Examples of the present invention are given below by way of illustration and not by way of limitation.

Examples 1 to 6 and Comparative Example 1

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Ingredients of the main component (A) which are shown in Table 1 were mixed uniformly at 60 °C for 5 minutes in mixing ratios which are also shown in Table 1. To component (A) were then added components (B) and (C), and the resulting mixture was stirred at 60 °C for 15 minutes using an agitator to obtain a resin composition for prepreg use. In this instance, each coating resin shown in Table 1 was used in an amount of 10% by weight based on the weight of diaminodiphenylsulfone used as the component (C).

Component (C) which was used in Examples 1 and 6 was prepared by the following method:

100 parts by weight of DDS particles having a size of from 10 to 30 μ m and 10 parts by weight of Nylon 12 particles having a size of 1 μ m were mixed in a high-speed mixer at a room temperature for 1 hour. The Nylon particles were adhered uniformly on the surface of the DDS particles electrostatically. The DDS particles having the Nylon 12 particles thereon were dispersed on a plastic sheet in such a manner that the DDS particles do not contact each other. Thus dispersed particles were placed in a thermostatic bath which was controlled to 170 °C and heated the particles for 30 minutes to melt and firmly adhere the Nylon 12 particles on the surface of the DDS particles.

Component (C) which was used in Examples 2 to 5 was prepared by the following method.

30 parts by weight of 3% by weight of an aqueous solution of a naphthalene sulfonic acid/formaldehyde condensation product and 120 parts by weight of 3% by weight of an aqueous solution of ethylene/maleic anhydride copolymer (average molecular weight: 80,000; as sodium salt) were mixed, and then DDS particles having particle sizes of 10 to 30 μ m were dispersed into the mixture. The pH of the dispersion was adjusted to 5.0. Then the mixture was further subjected to mixing using a homomixer at 10,000 rpm.

A melamine/formaldehyde primary polycondensation was prepared using 20 parts by weight of 37% formalin and 8 parts by weight of melamine. The primary polycondensation product was added to the above-described dispersion. The thus obtained mixture was then subjected to proceed a polycondensation reaction at 60 °C for 3 hours while stirring at 250 rpm.

The pH of the thus obtain d r action mixture was adjusted to 4.0 and th⁻ reaction was further continued at 80 °C for 2 hours. Particles coated with cur d melamine r sin thus obtained were recover d by filtering and washed with isopropyl alcohol. After then the particles were dried by air at 100 °C for 60 minutes to obtain component (C).

A resin film having a 30 μ m thickness was prepared from the thus obtained resin composition using a film coater at 70 °C. Carbon fibers (BESFIGHT IM-600 manufactur d by Toho Rayon Co., Ltd.; tensile strength: 600 kgf/mm²; modulus of elasticity: 30 \times 10³ kgf/mm²; diameter: 5 μ m) wer arranged in one direction on the thus prepared resin film, and impr gnation was effected continuously by hot-roll r at 80 °C under a pressure of 1 kg/cm² and at a speed of 3 m/min. In this way, a unidirectional prepreg containing carbon fiber in an amount of 145 g/m² and a resin content of 34% by weight was obtained.

A predetermined number of sheets were cut out from the thus obtained prepreg, 32 sheets or 8 sheets were laminated to one another and then subjected to autoclave molding at 180 °C under 5 kg/cm² for 2 hours to obtain a heat-cured molded plate. Thereafter, test pieces were cut out from the molded plate in order to measure the glass transition temperature thereof, compressive strength after 1500 in-lb/in impact (using the test piece compound of 32 sheets of prepreg), 0 ° compressive strength (compression strength in the same direction as that of the fibers - using the 8 sheets prepreg) under moist and warm conditions (tested at 82 °C after immersing the test piece in warm water of 71 °C for 2 weeks), and the gel time (at 180 °C) of the prepreg just after its preparation and after 2 months of storage at room temperature. The results are shown in Table 1.

In Comparative Example 1, a prepreg was prepared and its molded product was checked for its physical properties in the same manner as described above, except that component (B) was not used, with the results also shown in Table 1.

COMPARATIVE EXAMPLES 2 TO 4

Unidirectional prepregs were prepared and their properties were checked in the same manner as described above, except that 4,4'-diaminodiphenylsulfone which was not coated with a coating resin was used as the component (C). The results are shown in Table 1.

COMPARATIVE EXAMPLE 5

A unidirectional prepreg was prepared and its properties were checked in the same manner as described above, encapusulated except that 4,4'-diaminodiphenylsulfone which was not coated with a coating resin was used as component (C) and a cured melamine resin was also used in the resin composition by uniformly dispersing thereto. The results are shown in Table 1.

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5		cample	+			20	30	ı	20		15	15			1	ı	27	ч	1
		ive E	7			20	30	ı	20		30	1	ı			t	32	ı	ı
10		Comparative Example	7			20	30	20	ı		30	1	ı		1	ı	32	l.	í
15			4			20	30	20	i		ı	1	i		1	35	1	ı	1
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	_1	ple	4			20	30	ı	20		20	ı	7.0		1 .	35	ı	1	1
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40						-720	•		lified bisp		ide	lfone			odiphenyl sted with	odiphenyl sted with	odiphenylsu	nyldimethyl	sin
45				Mixing ratio	nponent (A)	Kaldite MY	Spikote 828	$\mathbb{E}_{\text{pikote}}$ 1001	Urethane modified bisphenol A resin	Component (B)	Polyetherimide	Polyethersulfone	Nylon 12	Component (C)	<pre>4,4'-Diaminodiphenyl sulfone (coated with nylon 12)</pre>	4,4'-Diaminodipheny sulfone (coated wit melamine)	4,4'-Diaminodiphenylsulfone	Dichlorophenyldimethyl urea	Melamine resin
50				Mixinç	CO	**	4	, **	2 4	CO	,		4	Co	~ W 12	~ 41 5	•	1	

	5	186 30	126 15 2	sheet	
5	Example 4	185	124	laminated	
10		188	130 17 3		
	Comparative 2 3	181	124 16 3	otopic f/mm ²)	
15		180	126 16 15	pseudo-isotopic	Tested at 82°C after immersion in 71°C hot water for 2 weeks the Measured at 180°C just after prepreg preparation. Measured at 180°C after 2 months of storage at room temperature. Mixing ratios in Table 1 are expressed by weight parts.
	9	181	123 16 15		z weer emperarts.
20	ru	200	135 19 16	ply	r ror n. room t ight p
25	Table 1 (cont'd) Example 2 3 4	170	110 17 15	ng 32	: wate iratio je at by wei
	Exc	185	128 10 6	t using	°C hot prepa storacessed
30	Table	188	125	g Calorimeter in-lb/in impact	82°C after immersion in 71°C hot water 101 2 wer at 180°C just after prepreg preparation. at 180°C after 2 months of storage at room temper ratios in Table 1 are expressed by weight parts.
	H	179	130 18 17	lorime lb/in	ersion fter p 2 mont
35		ה ה א	т *	ng Ca in-]	r inm just a ifter Table
40		cal properties Glass transition temp. °C, by DSC*1	<pre>impact *2 0° Compressive strength *3 Gel time *4 (min) Gel time *5 (min)</pre>	Scannin 1500	Cafte 80°C; 80°C; ios in
	•	ies lition l	<pre>impact *? impact *? 0° Compressive st Gel time *4 (min) Gel time *5 (min)</pre>	: Differentlal S Tested after (kgf/æm²).	at 82° dat 1 dat 1 ng rati
45		ropert trans y DSC	impact *2 0° Compressiv Gel time *4 (r Gel time *5	iffere ested kgf/æm	Tested at Measured a Measured a . Mixing
		Physical properties Glass transiti °C, by DSC*1 Compressive st	impac 0° Cc Gel t	Note (1): *1; Differentlal Scanning Calorimeter *2; Tested after 1500 in-lb/in im (kgf/mm²).	<pre>*3; Tested at *4; Measured *5; Measured Note (2): Mixing</pre>
50		Phys		Note	Note

On the basis of these results, it was confirmed that the prepregs obtained in Examples 1 to 6 had longer shelf-lives and the resulting composites had excellent thermal and mechanical characteristics and impact strength, as will as their longer shelf-lives, in comparison with those obtained in the Comparative Examples.

The resin composition of the present invention is capabl of supplying a composit with xcell nt impact str ngth without d teriorating other mechanical characteristics, including thermal r sistanc, and it has an xt nd d shelf-life in spite of th use of a diaminodiphenylsulfone as a curing agent.

Claims 5

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- 1. A resin composition which comprises:
 - (A) an epoxy resin,
 - (B) a thermoplastic resin selected from the group consisting of polyether imide, polyether sulfone, polysulfone, polycarbonate, polyetherether ketone, polyamide, and a mixture of two or more of
 - (C) particles of a diaminodiphenylsulfone compound whose cross-linking reactivity with the epoxy resin is prevented until the resin composition is subjected to molding or curing (i) by resin particles adhered on the surface of the diaminodiphenylsulfone particles or (ii) by a resin film coated thereon; the resin forming said resin particles or resin film being a polyamide, a modified urea resin, a cured melamine resin, a polyolefin, a paraffin, a modified paraffin or a mixture of two or more of these.
- 2. The resin composition as claimed in claim 1, wherein the epoxy resin is selected from the group consisting of glycidylamine epoxy resins, bisphenol epoxy resins, novolak epoxy resins, urethanemodified bisphenol A epoxy resins, alicyclic epoxy resins, and a mixture of two or more of these. 20
 - 3. The resin composition as claimed in claim 1, wherein the amount of component (B) is from 5 to 40 parts by weight per 100 parts by weight of component (A).
- The resin composition as claimed in claim 1, wherein the diaminodiphenylsulfone compound is 3,3'diaminodiphenyl sulfone, 3,4'-diaminodiphenyl sulfone or 4,4'-diaminodiphenyl sulfone.
 - 5. The resin composition as claimed in claim 1, wherein the amount of the diaminodiphenylsulfone compound is used in an epoxy equivalent/amine equivalent ratio of 1/(0.6 to 1.3).
 - The resin composition as claimed in claim 1, wherein the amount of the coating agent is from 5 to 20% by weight based on the weight of the diaminodiphenylsulfone compound.
- The resin composition as claimed in claim 1, wherein the coating resin coating melts or is destroyed at from 90 to 200 °C, under a pressure of from 1 to 5 kgf/cm², or at from 90 to 200 °C and under pressure 35 of from 1 to 5 kgf/cm².
 - The resin composition as claimed in claim 1, wherein the diaminodiohenylsulfone compound particle has a size of from 1 to 50 µm.
 - The resin composition as claimed in claim 1, wherein the composition contains a fiber reinforcement.
 - 10. The resin composition as claimed in claim 9, wherein the fiber reinforcement is selected from the group consisting of carbon fiber, glass fiber and aromatic polyamide fiber.
- 45 11. The resin composition as claimed in claim 9, wherein the amount of the fiber reinforcement is from 5 to 70% by volume based on the volume of the composition.
- 12. The resin composition as claimed in claim 1, wherein the resin particles are adhered onto the diaminodiphenylsulfone particles (i) separately from each other; (ii) to form a network structure; or (iii) to form continuous, non-continuous, porous or non-porous layer.
 - 13. A method for producing a resin composition which comprises mixing:
 - (A) an epoxy resin,
 - (B) a th rmoplastic resin sel cted from th group consisting of poly ther imide, polyether sulfone, polysulfone, polycarbonate, polyetherether ketone, polyamide, and a mixture of two or more of these; and

- (C) particl s of a diaminodiphenylsulfone compound whose cross-linking r activity with the poxy resin is pr v nted until th r sin composition is subj cted to molding or curing (i) by r sin particl s adh red on the surface of the diaminodiph nylsulfone particles or (ii) by a resin film coated thereon; the r sin forming said resin particles or r sin film being a polyamide, a modifi d urea r sin, a cur d melamine resin, a polyolefin, a paraffin, a modified paraffin or a mixture of two or more of these.
- 14. A method for producing a resin composition as claimed in claim 13, wherein a fiber reinforcement is impregnated with a mixture of components (A), (B) and (C) by a hot-melt method.
- 15. A method for producing a resin composition as claimed in claim 14, wherein the mixture is in the form of a film and the impregnation is conducted by hot-pressing.

Patentansprüche

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- 1. Harzzusammensetzung, die 15
 - (A) ein Epoxyharz,
 - (B) ein thermoplastisches Harz, das aus der aus Polyetherimid, Polyethersulfon, Polysulfon, Polycarbonat, Polyetheretherketon, Polyamid und einer Mischung aus zwei oder mehreren der genannten bestehenden Gruppen ausgewählt ist, und
- (C) Teilchen aus einer Diaminodiphenylsulfon-Verbindung, deren vernetzende Reaktivität zu dem 20 Epoxyharz (i) durch an der Oberfläche der Diaminodiphenylsulfonteilchen haftende Harzteilchen oder (ii) durch einen Harzfilmüberzug an diesen verhindert wird, bis die Harzzusammensetzung einem Formen oder Härten unterworfen wird; wobei das Harz, das die Harzteilchen oder den Harzfilm bildet, ein Polyamid, ein modifiziertes Harnstoffharz, ein gehärtetes Melaminharz, ein Polyolefin, ein Paraffin, ein modifiziertes Paraffin oder eine Mischung aus zwei oder mehreren der genannten ist, 25 enthält.
- Harzzusammensetzung nach Anspruch 1, in der das Epoxyharz aus der aus Glycidylamin-Epoxyharzen, Bisphenol-Epoxyharzen, Novolak-Epoxyharzen, Urethanmodifizierten Bisphenol A-Epoxyharzen, alicyclischen Epoxyharzen und Mischungen von zwei oder mehreren der genannten bestehenden Gruppe 30 ausgewählt ist.
 - Harzzusammensetzung nach Anspruch 1, in der die Menge der Komponente (B) 5 bis 40 Gewichtsteile pro 100 Gewichtsteile der Komponente (A) beträgt.
 - Harzzusammensetzung nach Anspruch 1, in der die Diamindiphenylsulfon-Verbindung 3,3'-Diamindiphenylsulfon, 3,4'-Diamindiphenylsulfon oder 4,4'-Diaminodiphenylsulfon ist.
- Harzzusammensetzung nach Anspruch 1, in der die Menge der Diaminodiphenylsulfon-Verbindung in einem Epoxyäquivalent/Aminäquivalent-Verhältnis von 1/(0,6 bis 1,3) eingesetzt wird. 40
 - Harzzusammensetzung nach Anspruch 1, in der die Menge des Überzugsagenzes 5 bis 20 Gew.%, bezogen auf das Gewicht der Diaminodiphenylsulfon-Verbindung, ist.
- 7. Harzzusammensetzung nach Anspruch 1, in der der beschichtende Harzüberzug unter einem Druck 45 von 1 bis 5 kgf/cm² zwischen 90 und 200 °C schmilzt oder zerstört wird.
 - Harzzusammensetzung nach Anspruch 1, in der das Teilchen der Diaminodiphenylsulfon-Verbindung eine Größe zwischen 1 und 50 µm hat.
 - Harzzusammensetzung nach Anspruch 1, wobei die Zusammensetzung eine Faserverstärkung enthält.
 - 10. Harzzusammensetzung nach Anspruch 9, in der die Faserverstärkung aus der aus Kohlenstoffaser, Glasfaser und Faser aus aromatischem Polyamid bestehenden Gruppe ausgewählt ist.
- 55 11. Harzzusammensetzung nach Anspruch 9, in der die Menge der Faserv rstärkung 5 bis 70 Vol%, bezogen auf das Volum in der Zusammensetzung, ist.

- 12. Harzzusammensetzung nach Anspruch 1, in der die Harzteilchen (i) getrennt voneinander; (ii) unter Bildung einer Netzstruktur; od r (iii) unt r Bildung ein r kontinui rlich n, nicht-kontinuierlich n, porös n oder nicht-porösen Schicht an den Diaminodiphenylsulfon-Teilchen haften.
- 13. Verfahren zur Herstellung einer Harzzusammensetzung, das ein Vermischen
 - (A) eines Epoxyharzes,
 - (B) eines thermoplastischen Harzes, das aus der aus Polyetherimid, Polyethersulfon, Polysulfon, Polycarbonat, Polyetheretherketon, Polyamid und einer Mischung aus zwei oder mehreren der genannten bestehenden Gruppen ausgewählt ist; und
 - (C) Teilchen aus einer Diaminodiphenylsulfon-Verbindung, deren vernetzende Reaktivität zu dem Epoxyharz (i) durch an der Oberfläche der Diaminodiphenylsulfonteilchen haftende Harzteilchen oder (ii) durch einen Harzfilmüberzug an diesen verhindert wird, bis die Harzzusammensetzung einem Formen oder Härten unterworfen wird; wobei das Harz, das die Harzteilchen oder den Harzfilm bildet, ein Polyamid, ein modifiziertes Harnstoffharz, ein gehärtetes Melaminharz, ein Polyolefin, ein Paraffin, ein modifiziertes Paraffin oder eine Mischung aus zwei oder mehreren der genannten ist; umfaßt.
- 14. Verfahren zur Herstellung einer Harzzusammensetzung nach Anspruch 13, in dem eine Faserverstärkung mit einem Gemisch der Komponenten (A), (B) und (C) nach einem Heißschmelzverfahren imprägniert wird. 20
 - 15. Verfahren zur Herstellung einer Harzzusammensetzung nach Anspruch 14, wobei das Gemisch in Form eines Film vorliegt und das Imprägnieren durch Heißpressen durchgeführt wird.

Revendications

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- 1. Composition de résine qui comprend :
 - (A) une résine époxy,
 - (B) une résine thermoplastique choisie dans le groupe constitué de polyétherimide, polyéthersulfone, polysulfone, polycarbonate, polyétheréthercétone, polyamide et un mélange de deux d'entre elles ou
 - (C) des particules d'un composé diaminodiphénylsulfone dont la réactivité de réticulation avec la résine époxy est masquée jusqu'à ce que la composition de résine est soumise au moulage ou au durcissement (i) par des particules de résine qui adhèrent à la surface des particules de diaminodiphénylsulfone ou (ii) par un film de résine qui est déposé sur elles ; la résine formant lesdites particules de résine ou le film de résine étant un polyamide, une résine urée modifiée, une résine mélamine durcie, une polyoléfine, une paraffine, une paraffine modifiée ou un mélange de deux ou plusieurs de ces produits.
- 2. Composition de résine selon la revendication 1, où la résine époxy est choisie dans le groupe constitué des résines époxy glycidylamine, résines époxy bisphénol, résines époxy novolaque, résines époxy bisphénol A uréthannemodifiées, et résines époxy alicycliques, et un mélange de deux ou plusieurs de ces résines.
- Composition de résine selon la revendication 1, où la quantité de composant (B) est de 5 à 40 parties en poids pour 100 parties en poids de composant (A).
 - Composition de résine selon la revendication 1, où le composé diaminodiphénylsulfone est 3,3'diaminodiphénylsulfone, 3,4'-diaminodiphénylsulfone ou 4,4'-diaminodiphénylsulfone.
 - Composition de résine selon la revendication 1, où la quantité de composé diaminodiphénylsulfone est utilisée en une proportion équivalent époxy/équivalent amine de 1/(0,6 à 1,3).
- Composition d résine selon la r v ndication 1, où la quantité de l'agent de revêtement est comprise entr 5 et 20 % n poids par rapport au poids du composé de diaminodiphénylsulfone. 55
 - 7. Composition de résine selon la revendication 1, où le r vêtement de résine de revêtement fond ou st détruit entre 90 et 200 °C, sous une pression de 1 à 5 kgf/cm², ou entre 90 et 200 °C et sous une

pression d 1 à 5 kgf/cm2.

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- Composition de résine selon la revendication 1, ou la particul de composé de diaminodiphénylsulfone a une taill de 1 à 50 μm.
- Composition de résine selon la revendication 1, où la composition contient un renfort de fibre.
- 10. Composition de résine selon la revendication 9, où le renfort de fibre est choisi dans le groupe constitué de fibre de carbone, fibre de verte et fibre de polyamide aromatique.
- 11. Composition de résine selon la revendication 9, où la quantité de fibre de renfort représente de 5 à 70 % en volume par rapport au volume de la composition.
- 12. Composition de résine selon la revendication 1, où les particules de résine adhérant aux particules de diaminodiphénylsulfone (i) séparées l'une de l'autre ; (ii) de façon à former une structure en filet ; ou 15 (iii) de façon à former une couche continue, non continue, poreuse ou non poreuse.
 - 13. Procédé de production d'une composition de résine qui comprend de mélanger :
 - (A) une résine époxy.
 - (B) une résine thermoplastique choisie dans le groupe constitué de polyétherimide, polyéthersulfone, polysulfone, polycarbonate, polyétheréthercétone, polyamide et un mélange de deux d'entre elles ou plus; et
 - (C) des particules d'un composé diaminodiphénylsulfone dont la réactivité de réticulation avec la résine époxy est masquée jusqu'à ce que la composition de résine est soumise au moulage ou au durcissement (i) par des particules de résine qui adhèrent à la surface des particules de diaminodiphénylsulfone ou (ii) par un film de résine qui est déposé sur elles ; la résine formant lesdites particules de résine ou le film de résine étant un polyamide, une résine urée modifiée, une résine mélamine durcie, une polyoléfine, une paraffine, une paraffine modifiée ou un mélange de deux ou plusieurs de ces produits.
 - 14. Procédé de production d'une composition de résine selon la revendication 13, où un renfort de fibre est imprégné d'un mélange de composants (A), (B) et (C) par un procédé de fusion à chaud.
- 15. Procédé de production d'une composition de résine selon la revendication 14, où le mélange est sous forme d'un film et on réalise l'imprégnation par pressage à chaud. 35

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